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(54) Title: DIEPISULFIDE BASED PREPOLYMERS AND THEIR USE IN THE OPTICAL FIELD

(57) Abstract: The room temperature polymerizable composition comprises at least one polymerizable monomer having at least one episulfide functionality and an effective amount of at least one (alkoxyphenyl) phosphine polymerization catalyst.

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DIEPISULFIDE BASED PREPOLYMERS AND THEIR USE IN THE OPTICAL FIELD

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(1) Field of the invention.

The present invention relates to stabilized, liquid diepisulfide based prepolymers, a method for preparing such prepolymers, a method for polymerizing curing the prepolymers into final polymer materials and their application for making transparent optical articles such as ophthalmic lenses.

(2) Description of the prior art

Diepisulfide based resins are of major interest in the optical field owing to their high refractive indexes of 1.7 or more.

Document US-A-5.807.975 discloses thermal polymerization and curing of diepisulfides, in particular for the manufacture of optical materials such as a lens material for spectacles. This thermal polymerization/curing of diepisulfides lasts for several hours, generally about 20 hours.

Document EP-0.942.027 discloses an optical resin prepared by curing a polymerizable composition comprising a (thio)epoxy compound having at least one intramolecular disulfide bond. The curing catalysts which may be used are typically tertiary amines, phosphines, Lewis acids, radical polymerization catalysts and cationic polymerization catalysts. Lengthy polymerization time are reported.

Document WO 00/26272 discloses a process for making thio containing resins which comprises polymerizing a composition comprising at least one diepisulfide compound and as polymerization catalyst, an alkaline metal, alkaline-earth metal or ammonium salt of an acid having a pKa ranging from 0.5 to 14.

Although, such a polymerization process leads to fast polymerization / curing of the composition, the process is very difficult to control due to the highly exothermic polymerization reaction and often results in a charred resin inappropriate for making optical articles such as ophthalmic lenses.

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SUMMARY OF THE INVENTION

It has now been discovered that it is possible to safely prepare a liquid diepisulfide based prepolymer which is stable in time, the prepolymer being able to be later safely polymerized / cured into a transparent resin useful for making optical articles such as ophthalmic lenses.

Thus, an object of the present invention is a stabilized, liquid, diepisulfide based prepolymer useful for making transparent final resins.

It is another object of the present invention to provide a process for making a stabilized, liquid, diepisulfide based prepolymer.

It is a further object of the present invention to provide a process for making a transparent optical article through fast polymerization / curing of a stabilized, liquid, diepisulfide based prepolymer.

These and other objects are attained by providing a stabilized, liquid, diepisulfide base prepolymer having a viscosity at 25°C ranging from 20 to 33000 cPs, preferably from 100 to 1000 cPs.

By a "stabilized" prepolymer there is meant a prepolymer the viscosity of which does not vary and particularly does not increase more than 20% when stored at 20°C over a period of 7 days, preferably 10 days and more preferably one month.

By a "liquid" prepolymer there is meant a prepolymer which can be poured and cast into a casting mold such as a casting mold for making ophthalmic lenses.

DETAILED DESCRIPTION OF THE INVENTION AND PREFERRED EMBODIMENTS

The stabilized, liquid, prepolymers of the invention are prepared by polymerizing a monomer composition comprising at least one diepisulfide monomer in the presence of an anionic initiator and a halogenated inhibitor.

Suitable diepisulfide monomers are diepisulfides of formula:

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in which R¹ and R² are, independently from each other, H, alkyl, aryl, alkoxy, aryloxy, alkythio or arylthio; R³ and R⁴, independently from each other,

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$$\begin{array}{c} \begin{array}{c} \begin{array}{c} \\ \\ \end{array} \\ \begin{array}{c} \\ \end{array} \end{array}$$

where R^a designates H, alkyl, aryl, alkoxy, aryloxy, alkythio or arylthio acid, n is an integer from 0 to 4 and m is an integer from 0 to 6.

R⁵, R⁶, R⁷, R⁸ each represents an hydrogen atom or a hydrocarbon group having 1 to 10 carbon atoms, and preferably a methyl group.

Preferably, each R⁵, R⁶, R⁷, R⁸ represents a hydrogen atom.

The preferred diepisulfide monomers are those having at least one intramolecular disulfide bond.

A preferred class of diepisulfide monomers is comprised of diepisulfide monomers of formula:

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in which R¹, R², R³ and R⁴ are defined as above.

In R^1 , R^2 , R^3 , R^4 and R^a , the alkyl and alkoxy groups are preferably C_1 - C_6 , more preferably C_1 - C_4 alkyl and alkoxy groups such as methyl, ethyl, propyl, butyl, methoxy, ethoxy, propoxy and butoxy.

Specific examples of diepisulfide monomers are bis(2,3-epithiopropyl)disulfide, bis(2,3-epithiopropyldithio)methane, bis(2,3-epithiopropyldithio)ethane, bis(6,7-epithio-3,4-dithioheptane)sulfide, 1,4-dithiomethyl)ethane, bis(6,7-epithio-3,4-dithioheptane)sulfide, 1,4-dithiomethyl)benzene, 1,6-bis(2,3-epithiopropyldithio)-2-(2,3-epithiopropyldithio)ethylthio)-4-thiahexane and 1,2,3-tris(2,3-epithiopropyldithio)propane.

The preferred diepisulfide monomer is bis(2,3-epithiopropyl)disulfide.

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$$H_2C$$
— CH — CH_2 — S — CH_2 — CH_2 — CH_2

The monomer compositions may comprise solely as polymerizable monomer a diepisulfide monomer of formula (I) or a mixture of such diepisulfide monomers or it may also include other polymerizable monomers.

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These additional monomers are compounds having two or more functional groups capable of reacting with an episulfide group or compounds having one or more of these functional groups and one or more of other homopolymerizable groups.

These additional monomers include epoxy compounds, polythiols, unsaturated ethylenic compounds such as vinyl ether or (meth)acrylate compounds, polyvalent carboxylic acids and anhydrides, thiocarboxylic acids, thio alcohols, thiophenols, polyphenols, amines and amides.

Preferred additional monomers are polythiol, epoxy compounds, acrylate and (meth)acrylate compounds, especially polythiol compounds.

Among the preferred epoxy compounds there may be cited the compounds of formulas:

$$H_{2}C \xrightarrow{R^{1}} C \xrightarrow{R^{b}} C \xrightarrow{R^{2}} CH_{2}$$
(III)

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$$H_{2}C \xrightarrow{\qquad \qquad } C \xrightarrow{\qquad } C \xrightarrow{\qquad \qquad } C \xrightarrow{\qquad } C \xrightarrow{\qquad \qquad } C \xrightarrow{\qquad } C \xrightarrow{\qquad \qquad } C \xrightarrow{\qquad } C \xrightarrow{\qquad \qquad } C \xrightarrow{\qquad } C \xrightarrow{\qquad \qquad } C \xrightarrow{\qquad \qquad } C \xrightarrow{\qquad } C \xrightarrow{\qquad \qquad } C \xrightarrow{\qquad } C \xrightarrow{\qquad \qquad } C \xrightarrow{\qquad \qquad } C \xrightarrow{\qquad \qquad } C \xrightarrow{\qquad } C \xrightarrow{\qquad } C \xrightarrow{\qquad } C \xrightarrow{\qquad } C$$

where X is O or S, R^b is H, alkyl or aryl and preferably alkyl or aryl, and R^1 , R^2 , R^3 , R^4 , m and n are defined as above.

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Typical examples of the unsaturated ethylenic compounds include ethylene glycol dimethacrylate, propylene glycol dimethacrylate, tetramethylene glycol dimethacrylate, polyethylene glycol dimethacrylate, trimethylolpropane triacrylate, pentaerythritolpropane trimethacrylate, ethoxylated bisphenol A diacrylate, 1,6-hexanediol dimethacrylate, urethane diacrylate, epoxy diacrylate, diallylphthalate, and divinyl benzene.

The polythiol monomer may be any suitable polythiol having two or more, preferably two to four, thiol functions.

The polythiol monomers can be represented by formula:

$$R'(SH)_{n'}$$
 (V)

In which n' is an integer from 2 to 6 and preferably 2 to 4, and R' is an organic group of valency equal to n'.

Useful polythiol monomers are those disclosed in EP-A-394.495 and US-A-4.775.733 and the polythiols corresponding to the following formulas:

$$\begin{array}{c} \text{HSCH}_2\text{ CH} \\ \text{OH} \end{array} \\ \begin{array}{c} \text{CH}_3 \\ \text{CH}_3 \end{array} \\ \begin{array}{c} \text{CH}_2\text{-CH} \\ \text{CH}_3 \end{array} \\ \begin{array}{c} \text{CH}_2\text{-CH} \\ \text{CH}_3 \end{array} \\ \end{array}$$

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$$\label{eq:hsc2} \text{HSC}_2\text{H}_4\text{OOC}_2\text{H}_4 = -\text{N} \\ \text{N} \\ \text{C} \\ \text$$

C₂H₅C(CH₂COOCH₂CH₂SH)₃

Among the preferred polythiol monomers there may be cited aliphatic mercaptoproprionate, 1tetrakis pentaerythritol such as polythiols 5 (1'mercaptoethylthio)-2,3-dimercaptopropane, 1-(2'-mercaptopropylthio)-2,3dimercaptopropane, 1-(-3'mercaptopropylthio)-2,3 dimercaptopropane, 1-(5' mercaptopentylthio)-1-(-4'mercaptobutylthio)-2,3dimercaptopropane, 2,3 dimercaptopropane, 1-(6'-mercaptohexylthio)-2,3-dimercaptopropane, 1,2bis(-4'-mercaptobutylthio)-3-mercaptopropane, 1,2-bis(-5' mercaptopentyl-10 1,2-bis(-6'-mercaptohexyl)-3-mercaptopropane, thio)-3-mercaptopropane, 1,2,3-tris(mercaptomethylthio)propane, 1,2,3-tris(-3'-mercaptopropylthio)propane, 1,2,3-tris(-2'-mercaptoethylthio)propane, 1,2,3-tris(-4'-mercaptobutylthio)propane, 1,2,3-tris(-6'-mercaptohexylthio)propane, 1,2-ethanedithiol, 1,1 propanedithiol, 1,2-propanedithiol, 1,3-propanedithiol, 15 2,2-propanedithiol, 1,6-hexanethiol-1,2,3-propanetrithiol, and 1,2-bis(-2'mercaptoethylthio)-3-mercaptopropane.

Among the preferred polythiols there may be cited polythiols of formulas:

HS—
$$CH_2$$
— CH_2

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mixtures thereof.

Examples of polythiol compounds include 1,2,3-trimethylolpropane tri(thioglycolate), pentaerythritol tetra(thioglycolate), pentaerythritol tetra(3-mercaptopropionate), 1,2,3-trimethylolpropane tri(3-mercaptoproprionate), thioglycerol, dithioglycerol, trithioglycerol, dipentaerythritol hexa(2-mercaptoacetate), and 3,4,5,6-tetrachloro-1,2-dimercapto benzene.

Among the (meth)acrylate compounds, there may be cited alkyl (C₁-C₆) (meth)acrylate such as methyl(meth)acrylate, ethyl(meth)acrylate, propyl(meth)acrylate and butyl(meth)acrylate.

The additional monomers usually represents 0 to 80% by weight preferably 0 to 50% by weight, of the total weight of polymerizable monomers present in the composition.

The composition may also include other optional ingredients classically used in the formulation of polymerizable compositions such as mold release agents, antioxidants, dyes and UV light absorbers. These and other ingredients may be employed in their customary amounts for their customary purposes.

The polymerizable monomer composition of the invention includes at least one anionic polymerizable initiator.

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Such anionic polymerization initiators are salts of formula:

$$M_{\mathbf{m}}^{p+} Y_{\mathbf{n}}^{-}$$
 (VI)

wherein M^{P+} is a cation selected from the group consisting of alkaline metals, alkaline earth metals, transitions metals and ammonium groups of formula NR⁺₄ in which R is an alkyl radical,

Y is an anion such as the corresponding acid YH has a pKa fulfilling the condition $0.5 \le pKa \le 14$,

p is the valency of the cation, and n = mxp.

The preferred metallic cation of the salts are Li⁺, Na⁺, K⁺, Rb⁺, Mg²⁺, Ca²⁺, Ba²⁺ and A1³⁺. The particularly preferred metallic cations are Li⁺, Na⁺, and K⁺ due to their absence of color and solubility in the composition. Transition metals are less preferred because the salts thereof lead to coloured compositions and therefore coloured polymerized resins.

The preferred NR_4^+ groups are those in which R is a C_1 - C_8 alkyl radical and more preferably, a methyl, ethyl, propyl, butyl or hexyl radical.

The salts shall be used in the polymerizable composition in an effective amount, i.e. an amount sufficient to promote the thermal or room temperature polymerization of the composition.

Generally, the salt will be present in amounts ranging, based on the total weight of the polymerizable monomers, from 5 to 10000 parts per million (ppm), preferably 100 to 5000 ppm.

Preferably, Y is an anion such as the corresponding acid YH which fulfils the condition $0.5 \le pKa \le 10$ and more preferably $0.5 \le pKa \le 8$.

Preferably, the anion Y is selected from the group consisting of thiocyanate, carboxylate, thiocarboxylate, acetylacetonate, diketone,

acetoacetic, ester, malonic, ester, cyanoacetic ester, ketonitrile and anion of formula RS wherein R is a substituted or non-substituted alkyl group or phenyl group.

Preferably, the alkyl group is a C₁-C₆ alkyl group, such as methyl, ethyl and propyl.

The preferred anions Y are SCN, acetylacetonate, acetate, thioacetate, formate and benzoate.

Examples of pKa (acid/base) corresponding to the polymerization initiators are given in the table below:

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ACID	ANION	Pka
HSCN	SCN	0.85
HCOOH	HCOO*	3.8
C ₆ H ₅ COOH	$C_6H_5COO^-$	4.2
· CH ₃ COOH	CH₃COO	4.8
C_6H_5SH	$C_6H_5S^-$	7.8
CH ₃ COCH ₂ COCH ₃	CH₃CO CHCOCH₃	9

Preferred polymerization initiators are KSCN and MeCOOK.

The polymerization initiator preferably also further includes at least one electro-donor compound.

The electro-donor compound is preferably a crown-ether.

The preferred crown-ethers may be represented by formula:

where R⁹ and R¹⁰ are polymethylene groups, preferably methylene or ethylene groups, and x is an integer from 3 to 6.

Preferred crown-ether is 1,4,7,10,13, 16-hexaoxacycloctadecane (18-crown-6).

The electro-donor compounds are present, based on total weight of the polymerizable monomers present in the composition in amounts ranging from 0 to 6%, preferably from 0.03 to 3% by weight.

Generally, the molar ratio $\frac{[electrodonor]}{[initiator]}$ ranges from 1 to 10.

The polymerization initiator is preferably introduced in the polymerizable composition as a solution in a solvent.

Appropriate solvents are alcohols, such as alcanols and mercaptoalcanols, ketones such as acetone, acetonitrile, dimethylsulfoxide and tetrahydrofuran. Preferred solvent is 2-mercaptoethanol.

The polymerizable composition also contains at least one halogenated polymerization inhibitor.

By polymerization inhibitor, it is meant a compound which decreases the rate of polymerization of a given polymerizable composition when compared to a similar polymerizable composition free of such polymerization inhibitor.

The halogenated inhibitor may be a chlorinated, a brominated, a iodinated or a mixed halogenated compound.

The halogenated inhibitor may be represented by the following formula:

$R(A)_y$ (VIII)

In which R is a carbon atom, or a hydrocarbon group such as an alkyl radical or an aromatic radical, A is an halogen or a halogeno alkyl radical and y is an integer from 1 to 4, preferably from 1 to 2.

The preferred halogenated inhibitors may be represented by formula:

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$$A_1$$
—(CH₂X) Z (IX)

where A₁ is an aryl radical eventually substituted, such as a phenyl, naphtyl, anthryl or biphenyl radical, X is a halogen atom such as I, Br or Cl, and z is a integer from 1 to 4, preferably 1 to 2.

Among the preferred halogenated inhibitors there may be cited:

CHa

$$CH_2CI$$
 CH_2CI CH_2CI CH_2CI

$$CH_2CI$$
 , OCN — CH_2CI

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OCN—CH
$$_2$$
CI , and CCl $_4$

The most preferred halogenated inhibitor is:

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The amount of halogenated inhibitor used may vary depending upon the monomers and the initiators used to formulate the polymerizable composition as well as the desired rate of polymerization. However, the amount of halogenated inhibitor shall be sufficient to avoid charring of the polymerizable composition.

Usually, the amount of inhibitor present in the polymerization composition expressed as the molar ratio of halogenated inhibitor to the polymerization initiator (with the exclusion of the electro-donor compound) will range from 0.1 to 350, preferably from 1 to 50, and more preferably from 1 to 10.

The monomer compositions may also include a variety of adjuvants such as photostabilizers, UV absorbents, antioxidants, anti-coloring agents, dyes, fillers and internal mold release agents in the usual proportions.

The prepolymer synthesis, i.e. polymerization of the monomer composition can be carried out in bulk, in solution, in suspension or in emulsion.

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Polymerization of the monomer compositions is preferably carried out in bulk at room temperature, by simply adding together the different ingredients of the compositions under stirring, then letting the polymerization proceed with or without stirring. Preferably the halogenated inhibitor is added to the monomer prior to the addition of the polymerization initiator.

As previously mentioned the prepolymers of the invention have a viscosity at 25°C ranging from 20 to 33000 centipoises, preferably 100 to 1000 centipoises.

All viscosity measurements are made with the same viscosimeter (Brookfield LVDVI) and following the same protocole but using S18 spindle for monomers and prepolymers of low viscosity (20 to 1000 cPs) and S31 spindle for prepolymers of higher viscosity (≥ 5000 and up to 50000 cPs).

Selection of spindle speed is made as follows:

For any given spindle/speed combination, the maximum range is equal to the spindle factor multiplied by 100 and the minimum range equals the factor multiplied by 10.

For the S-18 spindle, sample size is 8.0 ml and spindle factor is 30/N where N is the spindle speed (rpms) and for S-31 spindle, sample size is 10.0 ml and the spindle factor 300/N.

Thus, using S-18 spindle at 2.5 rpms the spindle factor will be 30/2.5 = 12 and the viscosity range 120 cPs to 1200 cPs.

Range efficiency (%) corresponds to the ratio of the measured viscosity for the sample divided by the maximum range and multiplied by 100. The higher the range efficiency, the more accurate is the viscosity measurement. Acceptable range efficiency is 10% to 99.9%.

Optimization of spindle speed may be effected as follows:

Assuming that at a spindle speed of 1.5 rpms the range efficiency is only 33.8%, doubling the spindle speed to 3.0 rpms would give a range efficiency of

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67.6%. Further increasing the spindle speed from 3.0 to 4.0 rpms would give a 25% increase of the range efficiency to 84.5%. But an increase of spindle speed from 4.0 to 5.0 rpms would give a 20% increase of the range efficiency to 101.4% which is an error (reading exceeds the 100% limit). Therefore, the viscosity measurements will be taken at a speed of 4 rpms with approximately a 84.5% range efficiency.

All viscosity measurements are made at 25°C.

All refractive index measurements are made at 25°C using sodium D line 589 nm.

The prepolymers of the invention can be safely cured into a transparent resin, in bulk polymerization.

Curing may be effected by simply heating the prepolymer or a mixture of the prepolymers at a temperature up to 80°C, preferably ranging from 40 to 80°C.

Curing can be conducted at room temperature by incorporating an additional amount of the polymerization initiator or another initiator to the prepolymer or mixture of prepolymers. The amount of additional or other polymerization initiator may vary depending upon the desired rate of curing but must be such that charring is avoided. This amount usually ranges from 100 to 2000 ppm.

Alternatively, initial raw monomers such as diepisulfides and/or polythiols as well as an additional amount of the polymerization initiator or another initiator can be added to the prepolymer or mixture of prepolymers, and the resulting composition can be cured at room temperature.

The amount of raw monomers that is usually added will range from 5 to 95%.

The additional amounts of polymerization initiator as previously indicated may be widely varied, but usually will range from 100 to 2000 ppm.

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Curing time up to the obtention of a gel usually ranges from 5 to 30 minutes.

Curing may be effected in a mold, such as a two parts mineral glass mold for obtaining an optical article such as an ophthalmic lens.

Whenever necessary, a mixing apparatus may be used for either the mixing step of the monomer components of the monomer composition leading to the prepolymer of the invention or the mixing steps of raw monomers or additives with the prepolymer of the invention.

Static mixers, small reactors with magnetic bar, mechanical mixers, screw mixers or ultrasonic mixing system may be used.

A screw mixer is preferably used.

In a preferred embodiment, the components to be mixed are added inside a small reactor chamber and then mixed with a screw mixer.

The following examples illustrate the present invention. In these examples, unless otherwise stated, all parts and percentage are by weight.

Examples 1 to 13 and comparative example C1

These examples illustrate the preparation of the prepolymers of the invention.

Several prepolymers were prepared by bulk polymerizing bis(2,3-epithiopropyl)disulfide using mixtures of MeCOOK and 18.crown.6 as the polymerization initiator in the presence of various amounts of benzylchloride as the inhibitor and eventually with various amounts of polythiols.

The polymerization initiator was added in the form of a solution in 2-mercaptoethanol.

Addition of the different ingredients and polymerization were made at room temperature. Magnetic stirring was used during the additions and stopped once the last ingredient added.

All polymerizations were done using 10g samples of bis(2,3-epithiopropyl)disulfide.

ta water a second

$$CH_2$$
 CH_2 CH_2 CH_2 CH_2 CH_2 CH_2

Compositions and results are given in Table I:

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TABLE I

Run N°	Polythiol A	MeCOOK	Molar ratio	Results
	(% by weight)	(ppm)	BzCl/MeCOOK	Time
1	-	52	0.142	Clear liquid after 6 days
2	-	43	0.331	Clear liquid after 6 days
3	-	61	0.422	Clear liquid after 6 days
4	-	51	1.004	Clear liquid after 6 days
5	-	49	1.572	Clear liquid after 6 days
· . 6	-	54	1.904	Clear liquid after 10 days
7	-	58	3.543	Clear liquid after 10 days
8	-	34	12.762	Clear liquid after 10 days
9	***	313	37.532	Clear liquid after 31 days
10	-	287	347.300	Clear liquid after 31 days
11	10.53	308	2.039	Slightly yellow viscous
				liquid after 37 days
12	10.18	300	5.324	Slightly viscous liquid
				After 5 days
13	10.98	321	8.843	Cloudy liquid after 37 days
C1	i -	304	0	Char after 50-86 min.

In runs 1 to 8 there was used 0.05-0.09% by weight of 2-mercaptoethanol (as solvent) and the molar ratio of 18-crown-6 to MeCOOK was 2.256.

In runs 9 to 13 and C1 there was used 0.5 to 0.6% by weight of 2-mercapto ethanol and the molar ratio of 18-crown-6 to the MeCOOK was 2.4.

In all runs there were added 0.4-0.5 ppm of Blue dye.

Polythiol A is a mixture of the following three polythiols in the respective weight percent amounts (molar amounts).

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Such polythiol A is described in US patent N°5.837.797 (example 1).

Blue dye: 1,4-[(2,6-dibromo-4-methylphenyl)amino]-1,4-

anthracenedione (also known as Morplas Violet 3B)

BzCl: Benzyl chloride

The liquid prepolymers made are perfectly miscible with the starting raw monomers.

Examples 14 and 15 and comparative example C'1

Prepolymers were prepared by bulk polymerizing bis(2,3-epithiopropyl)disulfide and polythiol A using a mixture of MeCOOK and 18-

crown-6 as initiator and CCl₄ as inhibitor in the presence of 2-ethylhexanoic acid.

The initiator was added in the form of a solution in 2-mercaptoethanol.

Compositions, polymerization conditions and results are given in Table

5 II.

Polymerizations were done using 10g samples of bis(2,3-epithiopropyl)disulfide.

TABLE II

Run N°	Molar ratio 2-	Polythiol A	CCl ₄	T(°C)	Stirred	Result	Time
	ethylhexanoic/	(%)	(%)				
	MeCOOK						
14	1.062	10.39	1.91	RT	No	Liquid	42 days
15	1.028	10.18	1.92	40°C	No	Liquid	42 days
Comparative	1.152	10.41	0	RT	No	Gel	1-16
C'1	-			•			hours

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The amount of MeCOOK was 295-325 ppm, the molar ratio 18-crown-6/Me COOK was 2.4 and the amount of 2-mercapto ethanol was 0.5-0.6%. RT = Room temperature (20°C).

Examples 16 to 26

Eleven prepolymers were prepared by polymerizing bis(2,3-epithiopropyl)disulfide using mixtures of MeCOOK and 18-crown-6 as initiator in the presence of different amounts of benzyl chloride as the inhibitor.

The polymerization initiator was added as a solution in 2-mercaptoethanol.

Addition of the different ingredients was made at room temperature under stirring. Polymerization was made at room temperature without stirring.

All polymerizations were done using 100g samples of bis(2,3-epithiopropyl)disulfide.

Viscosities of the prepolymers were measured as previously described. Compositions and results are given in Table III.

TABLE III

Run N°	Polythiol A	MeCOOK	Molar ratio		Viscosity (cPs)	
•	(% by weight)	(ppm)	BzCl/MeCOOK			,
16	•	300	2.100	245 (1 day)	298 (4 days)	299 (7 days)
17	-	302	4.091	59 (1 day)	73 (4 days)	74 (7 days)
18	-	300	6.264	35 (1 day)	44 (4 days)	44 (7 days)
19	5.057	303	2.127	91 (1 day)	167 (4 days)	175 (7 days)
20	5.052	301	4.079	37 (1 day)	49 (4 days)	51 (7 days)
21	5.052	302	5.815	35 (1 day)	40 (4 days)	41 (7 days)
22	10.009	301	2.099	253 (1 day)	323 (5 days)	320 (8 days)
23	10.020	301	4.059	47 (1 day)	53 (5 days)	54 (8 days)
24	10.036	301	6.066	37 (1 day)	39 (5 days)	40 (8 days)
25	•	51	0.526	51 (3 days)	49 (6 days)	49 (10 days)
26	•	48	0.139	60 (1 day)	64 (3 days)	67 (7 days)

In runs 16-24, there were used 0.5% by weight of 2-mercaptoethanol and 0.5 ppm of Blue dye. The molar ratio of 18-crown-6 to MeCOOK was 2.4. There was no stirring during the polymerization.

In runs 25 and 26 there were used 0.1% by weight of 2-mercaptoethanol and 0.5 ppm of Blue dye. Molar ratio of BzCl to Me COOK was 2.3 and the mixture was stirred during the polymerization.

The results show that the viscosity of the prepolymer increases with time to stabilize within 4 days. The final viscosity decreases with increasing amounts of inhibitor.

Examples 27 to 29

These examples illustrate the influence of the stirring and of the polymerization temperature on the resulting prepolymers.

Three prepolymers were prepared by polymerizing bis(2,3-epithiopropyl)disulfide using mixtures of MeCOOK and 18-crown-6 as initiator in the presence of different amount of benzylchloride as the inhibitor.

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The polymerization initiator was added as a solution in 2-mercaptoethanol.

Polymerization was effected either at room temperature or above with or without stirring.

All polymerizations were done using 100g samples of bis(2,3-epithiopropyl)disulfide.

Viscosities of the prepolymers were measured as previously described. Compositions and results are given in table (IV).

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TABLE IV

Run N°	MeCOOK (ppm)	Molar ratio 18-crown-6 / MeCOOK	Molar ratio BzCl/MeCOOK	2-mercapto- ethanol (%)	Stirring	T(°C)		Viscosity	
27	309	2.205	2.124	0.5	Yes	20	260 (1day)	281 (2 days)	282 (7 days)
28	300	2.400	2.100	0.5	No	20	245 (1 day)	248 (4 days)	299 (7 days);
29	308-309	2.205	2.058	0.5	Yes	40	1152 (1 day)	1291 (2 days)	1830 (7 days)

These examples show that stirring increases the rate of obtention of a stable prepolymer. Increasing the temperature increases the viscosity which keeps increasing with time without reaching a stable value.

Examples 30 to 34

These examples show the influence of the order of addition of the different reactants.

Several runs were carried out on 100g batches of bis(2,3-epithiopropyl)disulfides using different orders of addition of the reactants. Compositions, order of addition of the reactants and results are given in Table V.

The results show that the reaction is best controlled when benzyl chloride is first added to the monomer, followed by the initiator (run 34). The mixing upon addition is also important. It is worth noticing the lower reactivity of the system where the initiator solution was first mixed with benzyl chloride prior to addition to the monomer (run 33). The differences in the final viscosity of runs 32, 31 and 33 are probably due to differences in the amounts of benzyl chloride used rather than differences in the order of addition of the reactants.

TABLE V

Run N°	BzCl	Order of addition	Tmax (°C)	Viscosity (cPs)
		1) 0.6215 BDS		
		2) 58.7193 g M		
30	3.001	3) 0.1210 g BzCl	49.7@15 min	2541@116 hrs
ı		4) 0.6999 g initiator		
		5) 40.66 g M (S)		
		1) 0.3117 g BDS		-
		2) 2.3932 g M (S)		
		3) 10.0004 g M (S)		<u>516@92</u> hrs
31	3.338	4) 0.6861 g initiator	31.7@40 min	<u>566@140</u> hrs
		5) 0.1319 BzCl		
		6) 87.89 g M (S)		
		1) 0.3104 g BDS		
		2) 12.3970 g M (S)	·	<u>521@20</u> hrs
32	3.243	3) 0.1271 g BzCl (S)	31.5@40 min	578@48hrs
		4) 0.6805 g M (S)		• • •
		5) 87.98 g M (S)		
		1) 0.3175 g BDS		
		2) 12.3946 g M (S)		<u>237@20</u> hrs
33	3.571	3) [0.6831 g initiator +	≈ <u>26@30</u> min	<u>263@48</u> hrs
		0.1405 g BzCl(S)] (S)		
<u></u>		4) 88.00 g M (S)		
		1) 0.3077 g BDS		
		2) 100.42 g M (S)		203@1.87 hrs
34	3.756	3) 0.1486 g BzCl (S)	35.5@45 min	418@28 hrs
		4) 0.6869 g MeCOOK (S)		

BDS: solution made from 0.0037 g Blue Dye + 22.9947 g bis(2,3-epithiopropyl)disulfide

Initiator: solution made from 0.3914 g MeCOOK + 2.3557 g (18,6) crown + 6.0181 g 2-

5 mercaptoethanol

S: Stirred

M: bis(2,3-epithiopropyl)disulfide.

Examples 35 to 38

Four prepolymers were synthesized at room temperature using 100g batches of bis(2,3-epithiopropyl)disulfide monomer and different amounts of benzyl chloride. They were characterized by their viscosity and refractive index.

Compositions and results are given in Table VI:

TABLE VI

Run ^{a)} No	BzCl/MeCOOK	Viscosity	Viscosity	n_{D}	$n_{ m D}$
		(cPs)	(cPs)	@ 4 days	@ 19 days
	×	@ 4 days	@ 19 days		
36	2.224	29400 ^{b)}	28020°)	1.6866 ^{b)}	1.6865 ^{c)}
37	4.191	257	257	1.6656	1.6658
38	6.400	101	102	1.6587	1.6590
39	8.160	74	71	1.6559	1.6562

- a) All runs contained: [MeCOOK] ≈ 305 ppm, [(18,6)] ≈ 1832 ppm, [2-mercaptoethanol] ≈ 0.47%
- b) At 5 days
- 10 c) At 18 days

The refractive index of the prepolymers at 25°C decreases when the ratio BzCl/MeCOOK used increases. The values of refractive index are about the same between 4 and 19 days, which indicates that the prepolymers made are stable.

The viscosity of the prepolymers at 25°C decreases when the ratio BzCl/MeCOOK used increases. The values of viscosity are about the same between 4 and 19 days, which indicates that the prepolymers made are stable.

Examples 39 to 47

A series of nine chlorinated aromatics was evaluated as inhibitors, including benzyl chloride. All of them gave stable and clear prepolymers, except α , α dichlorotoluene that gave a char.

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In all runs, there were used 300 ppm MeCOOK, 1800 ppm of 18-crown-6 and 0.45% of 2-mercaptoethanol.

Different amounts of inhibitors were used as indicated by the molar ratios inhibitor/MeCOOK of Table VII. Results are also given in Table VII.

TABLE VII

Run N°	Inhibitor	Molar ratio	Experimental	Calculated
		Inhibitor/MeCOOK	Refractive index	viscosity (cPs)
			@ 7 days	
39	Benzyl chloride	2.019	1.6868	30070
	·	4.153	1.6659	285
		7.634	1.6567	76
40	1,2-bis(chloromethyl)	2.934	1.6546	59
	benzene	4.037	1.6522	44
		8.891	1.6478	28
41	1,3-bis(chloromethyl)	2.134	1.6760	1928
	benzene	4.616	1.6588	99
	· •	7.756	1.6528	48
42	1,4-bis(chloromethyl)	1.829	1.6708	673
	benzene	3.648	1.6607	129
	·	8.539	1.6516	42
- 43	4,4-	2.122	1.6676	378
	bis(chloromethyl)1,1-	4.506	1.6559	68
	biphenyl	7.503	1.6474	27
44	2-chloromethyl phenyl	1.836	1.6450	22
	isocyanate	4.030	1.6446	.21
		8.742	1.6447	21
45	4-chloromethyl phenyl	2.363	1.6472	27
	isocyanate	4.368	1.6454	22
		7.383	1.6453	22
46	1-chloromethyl 2,4-	2.821	1.6447	21
	diisocyanatobenzene	4.512	1.6449	21
		7.688	1.6454	22
47	α, α dichlorotoluene	3.052	_	
	•	3.272	-	-
·		8.537	. —	-
	and the state of the			

Calculated viscosities were determined using the graph of figure 1 which is a graph of Ln(viscosity) as a function of the refractive index.

Examples 48 to 51

These examples illustrate the use of the prepolymers of the invention for making cured transparent resins.

Samples of 10 g were prepared and placed in glass vials by adding to a prepolymer of the invention various amounts of polythiol A and MeCOOK. The prepolymers used contain 99.24% by weight bis(2,3-epithiopropyl)disulfide, 308 ppm MeCOOK, 1827 ppm 18-crown-6, 0.47% by weight 2-mercaptoethanol, 820 ppm of benzyl chloride and 0.5 ppm of Blue dye.

The mixture was stirred magnetically for about 10 s, then kept at room temperature with no stirring.

The amounts of reactants and the results are given in Table VIII.

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TABLE VIII

Run N°	Prepolymer	Polythiol A	Added [MeCOOK]	Result	Gel Time
	(%)	(%)	(ppm)		
48	88.7	10.4	77	Gel	> 24 hrs
49	88.3	10.3	307	Gel	< 24 hrs
50	88.2	10.2	355	Gel	< 24 hrs
51	88.0	10.4	405	Yellow gel	8 min

Examples 52 to 54

A mixture was prepared using the same prepolymer as in examples 47 to 50 and adding to this polymer, polythiol A and MeCOOK. The amounts of the reactants are indicated in Table IX.

This mixture was used to form 2 mm thick flat lenses having a diameter of 85 mm using a classical two parts mineral glass mold.

The lenses were cured 1 hour at 80°C. The resulting lenses are perfectly clear.

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TABLE IX

Run N°	Prepolymer	Polythiol A	Added [MeCOOK]	Gel Time
	(%)	(%)	(ppm)	(min)
52	88.7	9.7	413	60
. 53	88.4	9.6	537	40
54	87.6	9.7	871	11

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CLAIMS

- 1. A stabilized, liquid, diepisulfide based prepolymer having a viscosity at 25°C ranging from 20 to 33000 centipoises.
- 2. The prepolymer of claim 1, wherein the viscosity at 25°C ranges from 100 to 1000 centipoises.
- 3. The prepolymer of claim 1 resulting from the polymerization in the presence of an anionic polymerization initiator and a halogenated polymerization inhibitor of a monomer composition comprising at least one diepisulfide monomer of formula:

in which R¹ and R² are, independently from each other, H, alkyl, aryl, alkoxy, aryloxy, alkythio or arylthio; R³ and R⁴, independently from each other represent:

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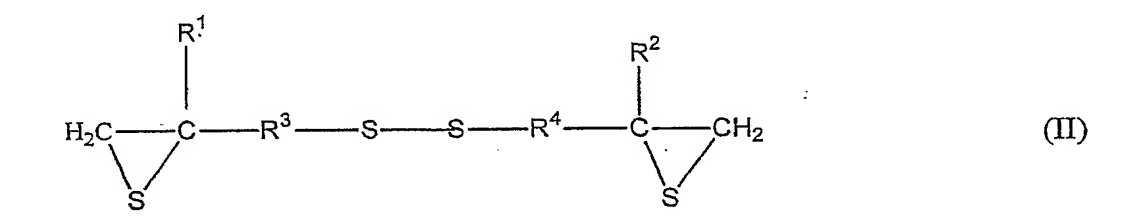
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where R^a designates H, alkyl, aryl, alkoxy, aryloxy, alkythio or arylthio acid, n is an integer from 0 to 4 and m is an integer from 0 to 6 and R^5 , R^6 , R^7 ,

R⁸ each represents a hydrogen atom or a hydrocarbon group having 1 to 10 carbon atoms.

4. The prepolymer of claim 3, wherein said at least one diepisulfide is a diepisulfide of formula:

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in which R¹, R², R³ are as defined in claim 3.

5. The prepolymer of claim 3, wherein said at least one diepisulfide is selected from bis(2,3-epithiopropyl)disulfide, bis(2,3-epithiopropyl-dithio)methane, bis(2,3-epithiopropyldithio)ethane, bis(6,7-epithio-3,4-dithioheptane)sulfide, 1,4-dithan-2,5-bis(2,3-epithiopropyldithiomethyl), 1,3-bis(2,3-epithiopropyl-dithiomethyl)benzene, 1,6-bis(2,3-epithiopropyldithio)-2-(2,3-epithiopropyl-dithioethylthio)-4-thiahexane and 1,2,3-tris(2,3-epithiopropyldithio)propage

epithiopropyldithio)propane.

- 6. The prepolymer of claim 3, wherein said at least one diepisulfide is bis(2,3-epthiopropyl)disulfide.
- 7. The prepolymer of claim 3, wherein the monomer composition further comprises at least one polythiol.
- 8. The prepolymer of claim 7, wherein said at least one polythiol is represented by formula:

$$R'(SH)_{n'}$$
 (V)

In which n' is an integer from 2 to 6 and preferably 2 to 4, and R' is an organic group of valency equal to n'.

9. The prepolymer of claim 8, wherein said at least one polythiol is selected from polythiols of formulas:

$$HS$$
—— CH_2 — C

ر

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mixtures thereof.

10. The prepolymer of claim 3, wherein the anionic polymerization initiator comprises a salt of formula:

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$$M_{\mathbf{m}}^{p+} Y_{\mathbf{n}}^{-}$$
 (VI)

wherein M^{P+} is a cation selected from the group consisting of alkaline metals, alkaline earth metals, transitions metals and ammonium groups of formula NR⁺₄ in which R is an alkyl radical,

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Y is an anion such as the corresponding acid YH has a pKa fulfilling the condition $0.5 \le pKa \le 14$,

p is the valency of the cation, and

n = mxp.

- 11. The prepolymer of claim 10, wherein the anionic polymerization initiator is MeCOOK or KSCN.
- 12. The prepolymer of claim 3, wherein the halogenated polymerization inhibitor comprises a compound of formula:

$$A_1$$
——(CH₂X) Z (IX)

where A_1 is an aryl radical eventually substituted, X is a halogen atom and z is an integer of 1 to 4.

13. The prepolymer of claim 3, wherein the halogenated polymerization inhibitor is selected from:

$$\begin{array}{c} \text{CH}_2\text{CI} \\ \\ \text{CH}_2\text{CI} \end{array}, \quad \text{CIH}_2\text{C} \\ \end{array}$$

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$$CH_2CI$$
 , OCN — CH_2CI OCN — CH_2CI , CH_2CI

- 14. The prepolymer of claim 13, wherein the halogenated polymerization inhibitor is benzyl chloride.
- 15. A process for making a stabilized, liquid, diepisulfide based prepolymer having a viscosity at 25°C ranging from 20 to 33000 centipoises which comprises polymerizing at room temperature a monomer composition comprising at least one diepisulfide monomer in the presence of at least one anionic polymerization initiator and at least one halogenated polymerization inhibitor.
- 16. The process of claim 15, wherein the halogenated polymerization inhibitor is added to the monomer composition prior to the addition of the anionic polymerization inhibitor.
- 17. The process of claim 15, wherein said at least one diepisulfide is a diepisulfide of formula:

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in which R^1 , R^2 , R^3 are as defined in claim 3.

18. The process of claim 15, wherein said at least one diepisulfide is selected from bis(2,3-epithiopropyl)disulfide, bis(2,3-epithiopropyl-dithio)methane, bis(2,3-epithiopropyldithio)ethane, bis(6,7-epithio-3,4-dithioheptane)sulfide, 1,4-dithan-2,5-bis(2,3-epithiopropyldithiomethyl), 1,3-bis(2,3-epithiopropyl-dithiomethyl)benzene, 1,6-bis(2,3-epithiopropyldithio)-2-

(2,3-epithiopropyl-dithioethylthio)-4-thiahexane and 1,2,3-tris(2,3-epithiopropyldithio)propane.

19. The process of claim 15, wherein said at least one diepisulfide is bis(2,3-epthiopropyl)disulfide.

20. The process of claim 15, wherein the monomer composition further comprises at least one polythiol.

21. The process of claim 15, wherein said at least one polythiol is represented by formula:

 $R'(SH)_{n'}$ (V)

In which n' is an integer from 2 to 6 and preferably 2 to 4, and R' is an organic group of valency equal to n'.

22. The process of claim 15, wherein said at least one polythiol is selected from polythiols of formulas:

$$_{\rm HS}$$
 — $_{\rm CH_2}$ — $_{$

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mixtures thereof.

23. The process of claim 15, wherein the anionic polymerization initiator comprises a salt of formula:

 M_{m}^{p+} Y_{n}^{-} (V)

wherein M^{P+} is a cation selected from the group consisting of alkaline metals, alkaline earth metals, transitions metals and ammonium groups of formula NR⁺₄ in which R is an alkyl radical,

Y is an anion such as the corresponding acid YH has a pKa fulfilling the condition $0.5 \le pKa \le 14$,

p is the valency of the cation, and

n = mxp:

- 24. The process of claim 15, wherein the anionic polymerization initiator is MeCOOK or KSCN.
 - 25. The process of claim 23, wherein the anionic polymerization initiator further comprises a crown-ether.
 - 26. The process of claim 25, wherein the crown-ether is 18-crown-6.

27. The process of claim 15, wherein the halogenated polymerization inhibitor comprises a compound of formula:

$$A_1$$
—(CH_2X) Z (IX)

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where A_1 is an aryl radical eventually substituted, X is a halogen atom and z is an integer of 1 to 4.

28. The process of claim 15, wherein the halogenated polymerization inhibitor is selected from:

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$$\begin{array}{c} \text{CH}_2\text{Cl} \ , \\ \text{ClH}_2\text{C} \end{array}$$

$$CH_2CI$$
 CH_2CI CH_2CI CH_2CI

$$-$$
CH $_2$ CI , OCN $-$ CH $_2$ CI

OCN—CH
$$_2$$
CI , and CCl $_4$

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- 29. The process of claim 15, wherein the halogenated polymerization inhibitor is benzyl chloride.
- 30. The process of claim 15, wherein the amount of anionic polymerization initiator present ranges from 5 to 10000 ppm based on the total weight of polymerizable monomers.
- 31. The process of claim 15, wherein the molar ratio of the halogenated inhibitor to the polymerization initiator ranges from 0.1 to 350.
- 32. A process for making a cured, transparent, diepisulfide based resin which comprises:
 - obtaining a stabilized, liquid, diepisulfide based prepolymer having a viscosity at 25°C ranging from 20 to 33000 centipoises;
 - adding to the prepolymer an effective amount of an anionic polymerization initiator; and
 - bulk polymerizing the prepolymer at least up to the obtention of a gel.
- 33. The process of claim 32, wherein bulk polymerization is carried out at room temperature.
- 34. The process of claim 33, wherein bulk polymerization up to the obtention of a gel lasts for 5 to 30 minutes.
- 35. The process of claim 32 which further comprises adding diepisulfide monomer and/or polythiol monomer prior to the bulk polymerization.
 - 36. The process of claim 35, wherein bulk polymerization is carried out at room temperature.
- 37. The process of claim 36, wherein bulk polymerization up to the obtention of a gel lasts for 5 to 30 minutes.
- 38. The process of claim 32, wherein bulk polymerization is carried out in a lens casting mold.
- 39. The process of claim 35, wherein bulk polymerization is carried out in a lens casting mold.
 - 40. Ophthalmic lens obtained by the process of claim 38.

- 41. Ophthalmic lens obtained by the process of claim 39.
- 42. A process for making a cured, transparent, diepisulfide based resin which comprises:
 - obtaining a stabilized, liquid, diepisulfide based prepolymer having a viscosity at 25°C ranging from 20 to 33000 centipoises;
 - adding to the prepolymer an effective amount of an anionic polymerization initiator;
 - bulk polymerizing the prepolymer with heating at least up to the obtention of a gel.

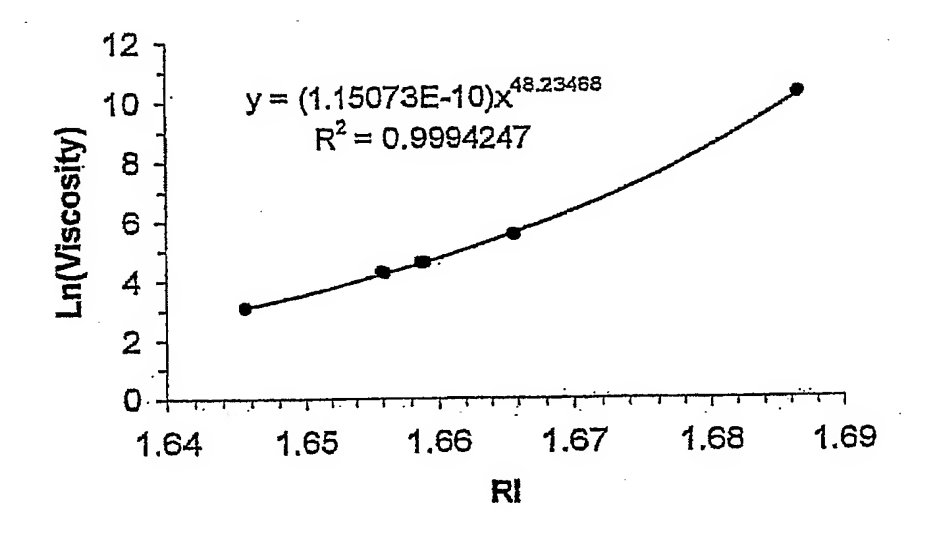


FIG. 1

INTERNATIONAL SEARCH REPORT

onal Application No Int

			P IP 01/15100
A. CLASSI IPC 7	FICATION OF SUBJECT MATTER C08G75/08 G02B1/04		·
According to	o International Patent Classification (IPC) or to both national class	sification and IPC	•
B. FIELDS	SEARCHED		
Minimum do IPC 7	cumentation searched (classification system followed by classification country of the country of	calion symbols)	
Documenta	tion searched other than minimum documentation to the extent th	at such documents are incli	uded in the fields searched
Electronic d	lata base consulted during the international search (name of data	a base and, where practical	, search lerms used)
EPO-In	ternal, WPI Data, PAJ		
C. DOCUM	ENTS CONSIDERED TO BE RELEVANT		
Category °	Citation of document, with indication, where appropriate, of the	e relevant passages	Relevant to claim No.
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X Furt	ther documents are listed in the continuation of box C.	χ Patent family	members are listed in annex.
A docum	ategories of cited documents: nent defining the general state of the art which is not dered to be of particular relevance	or priority date an	plished after the international filing date and not in conflict with the application but and the principle or theory underlying the
filing (document but published on or after the international date ent which may throw doubts on priority claim(s) or is cited to establish the publication date of another	involve an inventi	rular relevance; the claimed invention ered novel or cannot be considered to ve step when the document is taken alone
citatio "O" docum	nent referring to an oral disclosure, use, exhibition or means	cannot be considered document is com	ular relevance; the claimed invention ered to involve an inventive step when the bined with one or more other such docubination being obvious to a person skilled
"P" docum latert	nent published prior to the international filing date but than the priority date claimed		r of the same patent family
Date of the	actual completion of the international search	Date of mailing of	the international search report
1	18 April 2002	07/05/2	2002
Name and	mailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2 NL – 2280 HV Rijswijk	Authorized officer	
-	Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	0'Sull	ivan, T

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